

(12) United States Patent

Humayun et al.

(54) TUNGSTEN DEPOSITION PROCESS USING GERMANIUM-CONTAINING REDUCING AGENT

(71) Applicant: Lam Research Corporation, Fremont, CA (US)

(72) Inventors: Raashina Humayun, Los Altos, CA (US); Sudha Manandhar, Fremont, CA (US); Michal Danek, Cupertino, CA

Assignee: Lam Research Corporation, Fremont,

CA (US)

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(58) Field of Classification Search

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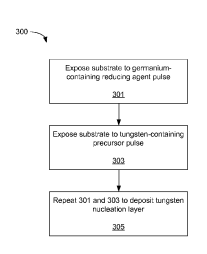
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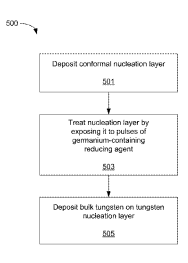
Primary Examiner — Caleb Henry (74) Attorney, Agent, or Firm — Weaver Austin Villenuve & Sampson LLP

(57)ABSTRACT

Methods for depositing low resistivity tungsten in features of substrates in semiconductor processing are disclosed herein. Methods involve using a germanium-containing reducing agent during tungsten nucleation layer deposition to achieve thin, low resistivity nucleation layers.

15 Claims, 11 Drawing Sheets





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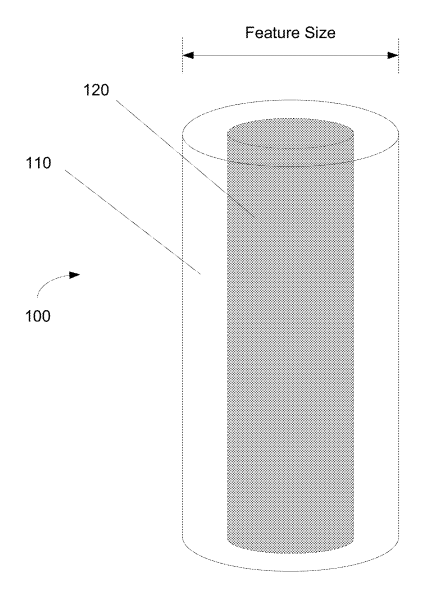
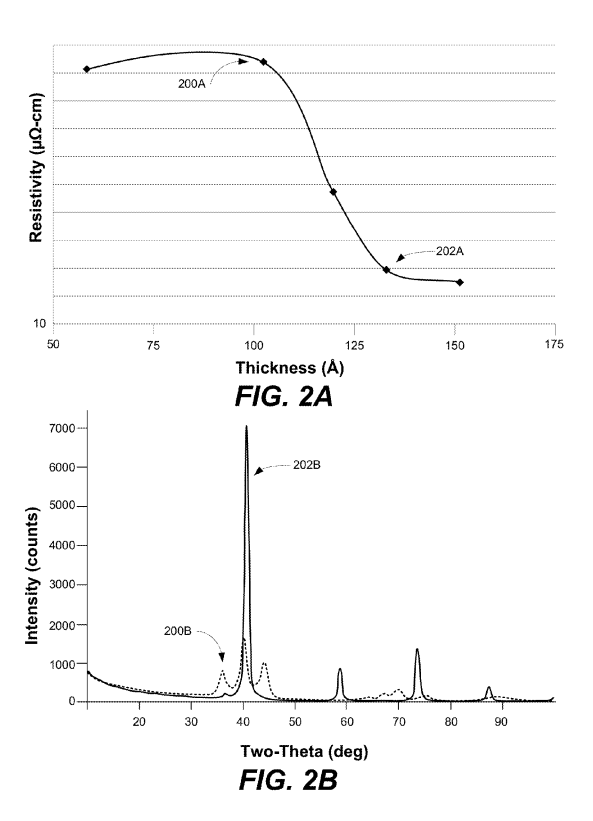


FIG. 1



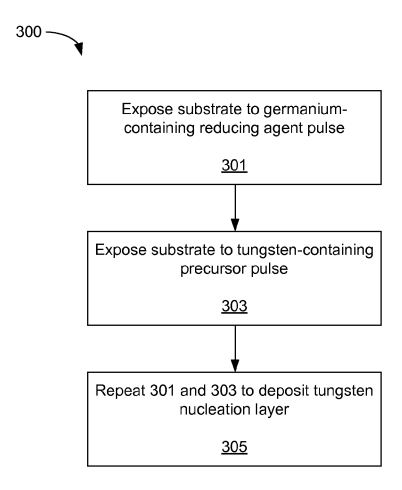


FIG. 3

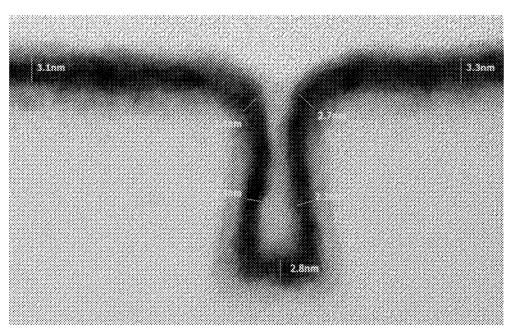


FIG. 4A

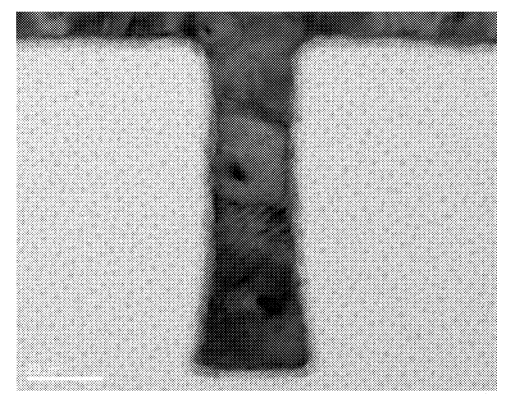


FIG. 4B

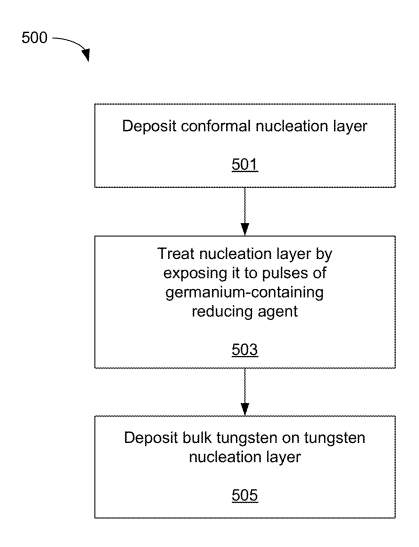


FIG. 5

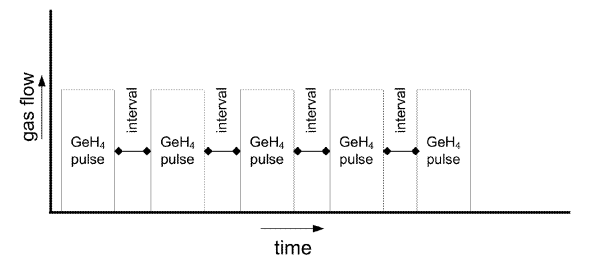


FIG. 6A

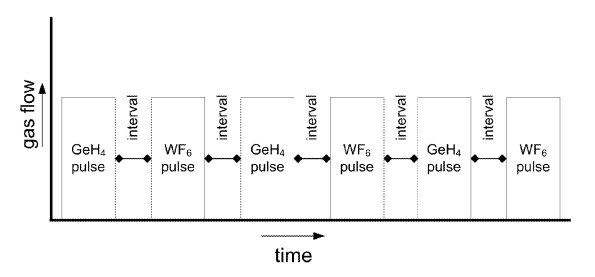


FIG. 6B

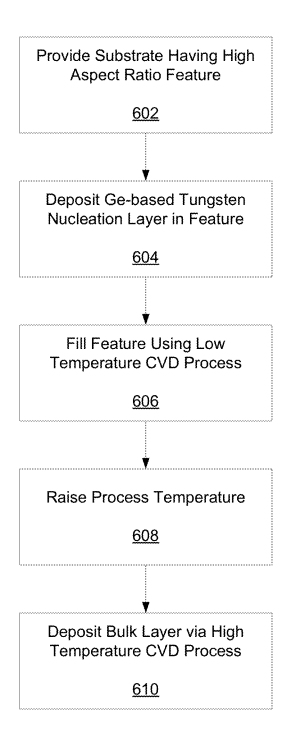
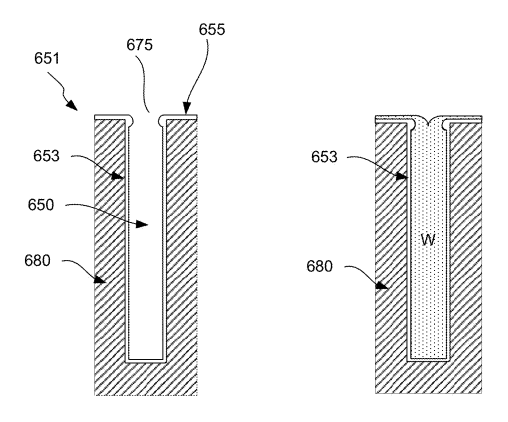


FIG. 6C



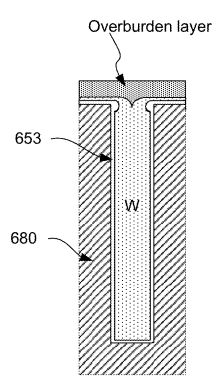


FIG. 6D

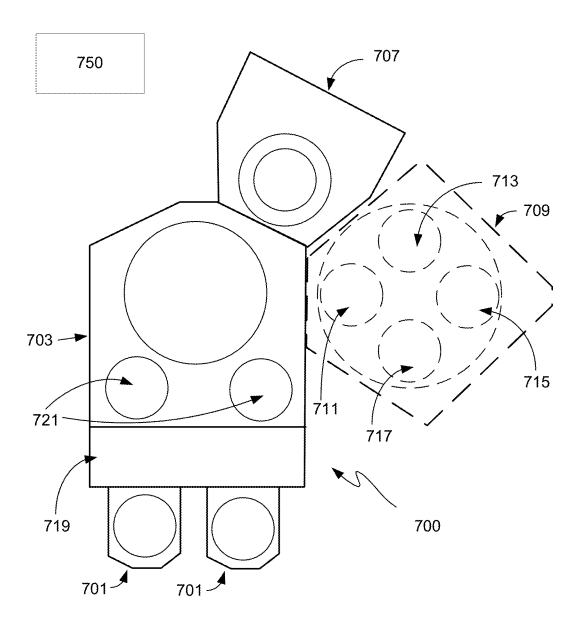
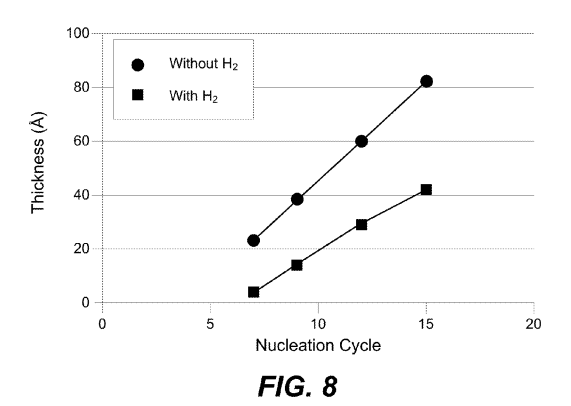
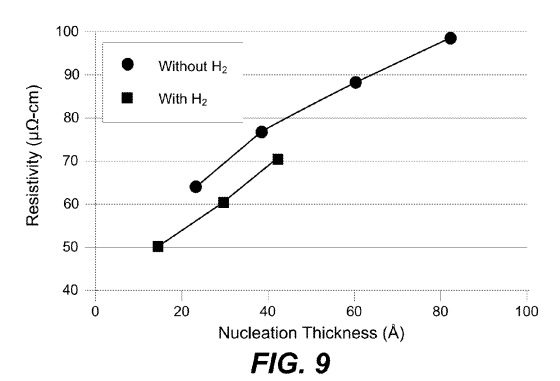
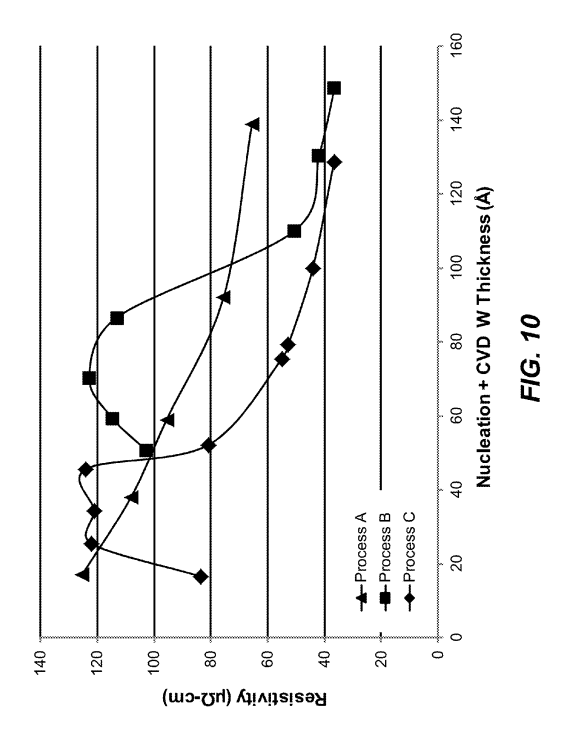


FIG. 7







TUNGSTEN DEPOSITION PROCESS USING GERMANIUM-CONTAINING REDUCING AGENT

CROSS-REFERENCE TO RELATED APPLICATION

This application claims benefit under 35 U.S.C. §119(e) of U.S. Provisional Patent Application No. 61/921,266, filed Dec. 27, 2013, and titled "TUNGSTEN NUCLEATION 10 PROCESS TO ENABLE LOW RESISTIVITY TUNGSTEN FEATURE FILL," which is herein incorporated by reference in its entirety and for all purposes. This application is also a continuation-in-part of U.S. patent application Ser. No. 14/097,160, filed Dec. 4, 2013, and titled "METHODS FOR DEPOSITING ULTRA THIN LOW RESISTIVITY TUNG-STEN FILM FOR SMALL CRITICAL DIMENSION CON-TACTS AND INTERCONNECTS," which is a continuation of U.S. patent application Ser. No. 12/755,259 (now U.S. Pat. No. 8,623,733), filed Apr. 6, 2010, and also titled METHODS $\,^{20}$ FOR DEPOSITING ULTRA THIN LOW RESISTIVITY TUNGSTEN FILM FOR SMALL CRITICAL DIMENSION CONTACTS AND INTERCONNECTS," which claims benefit under 35 U.S.C. §119(e) of U.S. Provisional Patent Application No. 61/169,954, filed Apr. 16, 2009 and titled 25 "METHOD FOR FORMING TUNGSTEN CONTACTS AND INTERCONNECTS WITH SMALL CRITICAL DIMENSIONS." Each of the prior applications is incorporated by reference herein in its entirety and for all purposes.

BACKGROUND

Deposition of tungsten-containing materials is an integral part of many semiconductor fabrication processes. These materials may be used for horizontal interconnects, vias 35 between adjacent metal layers, contacts between first metal layers and devices on the silicon substrate, and high aspect ratio features. In a conventional tungsten deposition process on a semiconductor substrate, the substrate is heated to the process temperature in a vacuum chamber, and a very thin 40 portion of tungsten film which serves as a seed or nucleation layer is deposited. Thereafter, the remainder of the tungsten film (the bulk layer) is deposited on the nucleation layer. The bulk layer is generally deposited more rapidly than the nucleation layer.

Increasingly thin tungsten electrical connects having very low resistance will enable smaller critical dimension devices. Although conventional methods are able to deposit nucleation layers, their ability to provide nucleation layers for the deposition of low resistivity tungsten in small critical dimension features is limited. For example, the formation of logic contacts has become more challenging as aspect ratios grow to more than 10:1. Void-free fill in aggressive features like these is problematic using conventional tungsten deposition techniques.

SUMMARY

Provided herein are methods for depositing tungsten, including methods for filling a feature on a substrate with 60 tungsten.

In one aspect, a method of filling a feature on a substrate with tungsten involves, prior to depositing a bulk tungsten layer, forming a tungsten nucleation layer by exposing the feature to alternating pulses of a germanium-containing 65 reducing agent and a tungsten-containing precursor. In various embodiments, the tungsten nucleation layer formed has a

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thickness between about 1 nm and 20 nm. In some embodiments, the tungsten nucleation layer formed has a thickness less than about 1 nm. In various embodiments, the amount of tungsten deposited per cycle is less than about 8 Å.

In certain embodiments, the tungsten nucleation layer is formed in a hydrogen atmosphere. In various embodiments, substantially all of the tungsten formed in the feature is alphatungsten. In many embodiments, the germanium-containing reducing agent is germane (GeH₄) or digermane (Ge₂H₆).

In some embodiments, the method also includes depositing a bulk tungsten layer by chemical vapor deposition. In certain embodiments, the grain growth during the bulk tungsten layer deposition extends from a first site where tungsten nucleated to a second site where tungsten nucleated in the feature. In many embodiments, substantially all of the tungsten formed during deposition of the bulk tungsten layer is alpha-tungsten.

Another aspect involves a method of filling a feature on a substrate with tungsten by forming a tungsten nucleation layer, and, after forming the tungsten nucleation layer and prior to depositing a tungsten bulk layer, treating the tungsten nucleation layer to pulses of a germanium-containing reducing agent. In various embodiments, there are no intervening pulses of other precursors during the treatment. In certain embodiments, intervening pulses of a tungsten-containing precursor are performed such that the amount of the tungstencontaining precursor pulsed during the treatment is less than the amount of the tungsten-containing precursor pulsed during the nucleation layer formation. In some embodiments, substantially no tungsten is deposited during the treatment with intervening pulses of tungsten-containing precursor. In many embodiments, the germanium-containing reducing agent is germane (GeH₄) or digermane (Ge₂H₆).

Another aspect involves an apparatus for filling a feature on a substrate including a multi-chamber apparatus with a chamber and a controller for controlling operations in the apparatus. The chamber may include a substrate support and one or more gas inlets configured to expose the substrate to gases. The controller may include machine-readable instructions for exposing the feature to alternating pulses of a germanium-containing reducing agent and a tungsten-containing precursor

In various embodiments, the controller also includes instructions for flowing hydrogen into the chamber while the feature is exposed to the alternating pulses of the germanium-containing reducing agent and the tungsten-containing precursor. In many embodiments, the germanium-containing reducing agent is germane (GeH₄) or digermane (Ge₂H₆). In various embodiments, the multi-chamber apparatus also includes a treatment chamber, and the controller also includes instructions for pulsing a germanium-containing reducing agent without intervening pulses of a tungsten-containing precursor.

These and other aspects are described further below with reference to the drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

 $FIG.\ 1$ is a schematic illustration of a feature filled with a tungsten nucleation and bulk layer according to certain embodiments.

FIGS. 2A and 2B are plots illustrating the presence of beta-tungsten versus alpha-tungsten in various embodiments.

FIG. 3 is a process flow diagram illustrating operations in a method of filling a feature with tungsten according to various embodiments.

FIG. **4**A is an example of an image of a feature with a deposited tungsten nucleation layer according to various embodiments.

FIG. **4**B is an example of an image of a feature with a deposited tungsten nucleation layer and tungsten bulk layer saccording to various embodiments.

FIG. 5 is a process flow diagram illustrating operations in a method of filling a feature with tungsten according to various embodiments.

FIGS. 6A and 6B illustrate gas pulse sequences in a low resistivity treatment according to various embodiments.

FIG. 6C is a process flow diagram illustrating operations in a method of filling a feature with tungsten according to various embodiments.

FIG. 6D is a schematic illustration of feature cross-sections at various stages of a process according to certain embodiments.

FIG. 7 is a schematic illustration of a processing system suitable for conducting tungsten deposition processes in 20 accordance with disclosed embodiments.

FIG. 8 is a plot illustrating film thickness as a function of nucleation deposition cycle in accordance with disclosed embodiments.

FIG. **9** is a plot illustrating resistivity as a function of film 25 thickness in accordance with disclosed embodiments.

FIG. 10 is a plot illustrating resistivity of deposited tungsten films in accordance with disclosed embodiments.

DETAILED DESCRIPTION

In the following description, numerous specific details are set forth to provide a thorough understanding of the presented embodiments. The disclosed embodiments may be practiced without some or all of these specific details. In other 35 instances, well-known process operations have not been described in detail to not unnecessarily obscure the disclosed embodiments. While the disclosed embodiments will be described in conjunction with the specific embodiments, it will be understood that it is not intended to limit the disclosed 40 embodiments.

There are various challenges in tungsten (W) fill as devices scale to smaller technology nodes. One challenge is preventing an increase in resistance due to the thinner films in contacts and vias. As features become smaller, the tungsten con- 45 tact or line resistance increases due to scattering effects in the thinner tungsten films. Low resistivity tungsten films minimize power losses and overheating in integrated circuit designs. Conventional chemical vapor deposition (CVD) tungsten deposition processes involve depositing a nucleation 50 layer followed by CVD bulk tungsten deposition. While efficient tungsten deposition processes use tungsten nucleation layers, these layers typically have higher electrical resistivities than the bulk layers. Barrier layers deposited in contacts, vias, and other features, may also have high resistivities. 55 Further, the thin barrier and tungsten nucleation films occupy a larger percentage of smaller features, increasing the overall resistance in the feature. Resistivity of a tungsten film depends on the thickness of the film deposited, increasing at very low thicknesses due to boundary effects.

FIG. 1 shows a volume occupied by a nucleation film 110 and a bulk tungsten material 120 in a via or contact structure 100 that may be formed on a substrate. Because the resistivity of the nucleation layer is higher than that of the bulk layer $(\rho_{nucleation} > \rho_{bulk})$, the thickness of the nucleation layer 65 should be minimized to keep the total resistance as low as possible. On the other hand, the tungsten nucleation should be

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sufficiently thick to fully cover the underlying substrate to support high quality bulk deposition.

Features, such as via or contact holes, may be characterized by one or more of narrow and/or re-entrant openings, constrictions within the feature, and high aspect ratios. One example of a feature can include a feature hole in a substrate or a layer on the substrate, such as the feature shown in FIG. 1. The substrate may be a silicon wafer, e.g., a 200-mm wafer, a 300-mm wafer, or a 450-mm wafer, including wafers having one or more layers of material, such as dielectric, conducting, or semi-conducting material deposited thereon. The feature may be formed in one or more of these layers. In some embodiments, the feature may have an aspect ratio of at least about 2:1, at least about 4:1, at least about 6:1, at least about 10:1, or higher. The feature hole may also have a dimension near the opening, e.g., an opening diameter or line width of between about 10 nm to 500 nm, for example between about 25 nm and about 300 nm. A feature hole may be referred to as an unfilled feature or a feature.

A feature that may have a re-entrant profile may have a profile that narrows from the bottom, closed end, or interior of the feature to the feature opening. In various embodiments, the feature may have an under-layer, such as a barrier layer or adhesion layer. Non-limiting examples of under-layers include dielectric layers and conducting layers, e.g., silicon oxides, silicon nitrides, silicon carbides, metal oxides, metal nitrides, metal carbides, and metal layers. In certain embodiments, the under-layer may be titanium nitride (TiN), titanium metal (Ti), tungsten nitride (WN), titanium aluminide (TiAl), or a titanium oxide (TiO_x).

Features may have one or more constrictions at a midpoint within the features such that pinch off may occur during deposition of tungsten and deposited tungsten blocks further deposition past the constriction before that portion of the feature is filled. Such features may be used in logic devices, such as in a word line in a vertical NAND (VNAND) structure.

For the purposes of this description, "near the opening" is defined as an approximate position or an area within the feature (i.e., along the sidewall of the feature) corresponding to between about 0% and about 10% of the feature depth measured from the field region. In certain embodiments, the area near the opening corresponds to the area at the opening. Further, "inside the feature" is defined as an approximate position or an area within the feature corresponding to between about 20% and about 60% of the feature depth measured from the field region on the top of the feature. Typically, when values for certain parameters (e.g., thicknesses) are specified "near the opening" or "inside the feature," these values represent a measurement or an average of multiple measurements taken within these positions/areas. In certain embodiments, an average thickness of an under-layer near the opening is at least about 10% greater than that inside the feature. In more specific embodiments, this difference may be at least about 25%, at least about 50%, or at least about 100%. Distribution of a material within a feature may also be characterized by its step coverage. For the purposes of this description, "step coverage" is defined as a ratio of two thicknesses, i.e., thickness of the material inside the feature 60 divided by the thickness of the material near the opening. In certain examples, the step coverage of the liner or other under-layer is less than about 100% or, more specifically, less than about 75% or even less than about 50%.

A tungsten nucleation layer may be deposited in a feature to conformally coat the sidewalls and bottom of the feature, or for the case of laterally oriented features having openings on both sides, the sidewalls of the feature. In general, a nucle-

ation layer is a thin conformal layer that serves to facilitate the subsequent formation of a bulk material thereon. Conforming to the underlying feature bottom and sidewalls can be critical to support high quality deposition. Nucleation layers are often deposited using atomic layer deposition (ALD) or pulsed 5 nucleation layer (PNL) methods.

In a PNL technique, pulses of reactant are sequentially injected and purged from the reaction chamber, typically by a pulse of a purge gas between reactants. A first reactant can be adsorbed onto the substrate, available to react with the next 10 reactant. The process is repeated in a cyclical fashion until the desired thickness is achieved. PNL is similar to ALD techniques. PNL is generally distinguished from ALD by its higher operating pressure range (greater than 1 Torr) and its higher growth rate per cycle (greater than 1 monolayer film 15 growth per cycle). Chamber pressure during PNL deposition may range from about 1 Torr to about 400 Torr. In the context of the description provided herein, PNL broadly embodies any cyclical process of sequentially adding reactants for reaction on a semiconductor substrate. Thus, the concept embod- 20 ies techniques conventionally referred to as ALD. In the context of the disclosed embodiments, CVD embodies processes in which reactants are together introduced to a reactor for a vapor-phase reaction. PNL and ALD processes are distinct from CVD processes and vice versa.

In many instances of tungsten nucleation layer deposition, the one of the sequentially-injected reactants may be a boroncontaining reducing agent (such as borane (BH₃) or diborane (B₂H₆)) or a silicon-containing reducing agent (such as silane (SiH₄)). However, both boron-containing reducing agents 30 and silicon-containing reducing agents present some concerns. Use of diborane (B₂H₆) as a reducing agent results in thicker nucleation film deposition per cycle. This is because diborane deposition is not self-limiting. Therefore, instead of depositing a saturated monolayer on the surface, diborane 35 may accumulate and grow on the surface of the feature, thereby yielding more diborane on the surface. The thicker layer of diborane results in a thicker nucleation layer of tungsten when the tungsten-containing precursor reacts with the diborane. Thus, when using diborane, tungsten nucleation 40 films are often deposited at a thickness of about 4 Å to about 12 Å per cycle.

In some cases, nucleation layer deposition is followed by a post-deposition treatment to improve resistivity, such as treating the nucleation layer with sequential pulses of a boroncontaining reducing agent such as diborane (B_2H_6). Another alternative is to expose the nucleation layer to alternate pulses of diborane and tungsten hexafluoride (WF₆). Although diborane-treated nucleation layers may have lower resistivity, potential issues still arise due to the presence of boron-10 ^{10}B), an isotope of boron, in the bulk tungsten film.

Presence of boron-10 causes integration issues, such as processing defects during chemical mechanical planarization (CMP), or soft error rate defects due to interaction of isotope boron-10 with thermal neutrons. Specifically, boron-10 reacts 55 with chemicals during CMP to form soluble boric acid, which leads to edge erosion, plug pull out, and other defects. These defects are of particular concern for deposited tungsten films in logic devices.

Another concern in using boron-containing reducing 60 agents may be its effect on the type of tungsten that forms in the feature. FIG. 2A is a plot depicting resistivity of tungsten films deposited using diborane and tungsten hexafluoride pulses for the nucleation layer deposition, followed by diborane pulses for post-deposition treatment, and lastly bulk 65 CVD tungsten deposition. Points 200A and 202A represent the resistivity of tungsten films of about 105 Å and 130 Å,

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respectively, deposited using this method with bulk CVD tungsten deposited at 300° C. The y-axis represents the resistivity of the overall deposited tungsten including both the nucleation layer and the bulk CVD tungsten, and the x-axis represents the thickness of the entire tungsten deposition including both the nucleation layer and the bulk CVD tungsten. As shown, there is a slight increase in the resistivity curve for boron-based tungsten film depositions prior to the resistivity decreasing with thickness. Note the lower resistivity of the tungsten film at point 202A. It should be noted that the resistivity curve characteristics may also depend on the CVD process employed; a boron-based nucleation layer followed by CVD at 395° C. did not show the same rise in resistivity as thickness increased.

The tungsten films represented by points 200A and 202A were evaluated using x-ray diffraction. The film represented by point 200A in FIG. 2A shows a peak 200B in FIG. 2B corresponding to presence of beta-tungsten (β -W). The film represented by point 202A shows a peak 202B corresponding to the high presence of alpha-tungsten (α -W). This data indicates that the presence of beta-tungsten increases the resistivity of the overall tungsten film. Beta-tungsten has a metastable A15 cubic crystalline structure and exhibits higher resistivity than the stable body-centered cubic crystalline structure of alpha-tungsten. Without being bound by a particular theory it is believed that boron-based nucleation layers may lead to the presence of higher resistivity beta-tungsten in tungsten films at certain thicknesses. As discussed further below, tungsten films deposited on top of germanium-based nucleation layers do not exhibit the resistivity behavior shown in FIG. 2A; rather the resistivity decreases with increasing thickness.

It is desirable for nucleation layers to form a good surface for bulk tungsten deposition. Silane (SiH₄) has been used in nucleation layer deposition as an alternative to boron-containing reducing agents. However, for silane-based nucleation layers to yield good bulk tungsten growth, a thicker layer of the tungsten nucleation layer is typically needed, such as a layer at least greater than 50 Å thick. Since the resistivity of a tungsten nucleation layer is higher, this results in an overall higher resistivity tungsten film. The lack of granularity in the thickness per cycle also results in reduced process control.

Provided herein are methods of filling features with tungsten and related systems and apparatuses. Examples of application include logic and memory contact fill, DRAM buried word line fill, vertically integrated memory gate/word line fill, and 3-D integration with through-silicon vias (TSVs). The methods described herein can be used to fill vertical features, such as in tungsten vias, and horizontal features, such as VNAND word lines. The methods may be used for conformal and bottom-up or inside-out fill.

It has been found that tungsten films deposited using germanium-containing reducing agents can provide low resistivity, thin tungsten nucleation films that yield alpha-tungsten growth, with the resulting films causing has little or no defects in logic devices. In particular, tungsten nucleation film deposition using germanium-containing reducing agents deposits thinner films per cycle, thereby lowering resistivity and providing finer control over the resulting thickness of the nucleation film. These thin nucleation layers also promote good tungsten growth for deposition of the tungsten bulk layer. Moreover, when a tungsten-containing precursor reacts with the germanium-containing deposited film in each cycle, the amount of germanium-containing deposited film left on the

substrate as measured by x-ray photoelectron spectroscopy is little, suggesting that most of the film in the feature is reduced to elemental tungsten.

FIG. 3 is a process flow diagram of a method for filling features with tungsten in accordance with various embodiments. In some embodiments, method 300 may be performed at a temperature between about 200° C. and about 400° C. In some embodiments, method 300 may be performed at a pressure between about 1 Torr and about 300 Torr. Temperatures and pressures outside these ranges may be used according to particular implementations. In operation 301, a substrate is exposed to a pulse of a germanium-containing reducing agent. As an example, the substrate may be a substrate one or more features to be filled with tungsten. According to various embodiments, the substrate feature has an aspect ratio of at least 10:1, at least 15:1, at least 20:1, at least 25:1 or at least 30:1. Also according to various embodiments, the feature size is characterized by the feature opening size in addition to or instead of the aspect ratio. The opening may be from 10 nm to 20 100 nm, or 10 nm to 50 nm wide. For example, in certain embodiments, the methods may be advantageously used with features having narrow openings, regardless of the aspect ratio. In certain embodiments, the recessed feature is formed within a dielectric layer on a substrate, with the bottom of the 25 feature providing contact to an underlying metal layer. Also in certain embodiments, the feature includes a liner/barrier layer on its sidewalls and/or bottom. Examples of liner layers include Ti/TiN, TiN and WN. In addition to or instead of diffusion barrier layers, the feature may include layers such as 30 an adhesion layer, a nucleation layer, a combination of thereof, or any other applicable material lining the sidewalls and bottom of the feature. In certain embodiments, the feature is a re-entrant feature; that is the liner layer or other material forms an overhang that partially blocks the feature opening. 35

Any germanium-containing compound that can decompose or react to form a layer capable of reducing the tungstencontaining precursor to form elemental tungsten may be used. Examples include germanes, such as $\text{Ge}_n H_{n+4}$, $\text{Ge}_n H_{n+6}$, $\text{Ge}_n H_{n+8}$, and $\text{Ge}_n H_m$, where n is an integer from 1 to 10, and 40 n is a different integer than m. Other germanium-containing compounds may also be used, e.g., alkyl germanes, alkyl germanium, aminogermanes, carbogermanes, and halogermanes. In general, halogermanes may not have significant reducing potential but there may be process conditions and 45 tungsten-containing precursors suitable for film formation using halogermanes.

Example pulse times can be between about 0.25 seconds and about 30 seconds, about 0.25 seconds to about 5 seconds, or about 0.5 seconds to about 3 seconds. The pulse may be 50 sufficient to saturate the surface of the features in the substrate. In some embodiments, the pulse may be sufficient to oversaturate the surface of the features in the substrate. In some embodiments, a carrier gas may be used, such as, for example, argon (Ar), helium (He), or nitrogen (N₂).

In various embodiments, operation 301 may be performed in a hydrogen environment. For example, hydrogen may be flowed to a chamber where the substrate is processed at a hydrogen-to-germanium-containing reducing agent ratio of at least 10:1, or at least 50:1, or at least 70:1, or at least 100:1. 60 As discussed further below, deposition in a hydrogen environment may decrease the thickness deposited per cycle, as well as decrease the resistivity of the deposited tungsten film.

After operation 301, there may be an optional purge step to purge excess germanium-containing reducing agent still in 65 gas phase that did not adsorb to the surface of the feature. A purge may be conducted by flowing an inert gas at a fixed

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pressure thereby reducing the pressure of the chamber and re-pressurizing the chamber before initiating another gas exposure

Next, in operation 303, the substrate is exposed to a tungsten-containing precursor pulse. The tungsten-containing precursor reacts with the deposited layer of germanium to form elemental tungsten. Examples of tungsten-containing precursors include tungsten hexafluoride (WF₆), tungsten hexachloride (WCl₆), or tungsten hexacarbonyl (W(CO)₆). In some embodiments, the tungsten-containing precursor is a halogen-containing compound, such as tungsten hexafluoride. Organo-metallic precursors and precursors that are free of fluorine, such as MDNOW (methylcyclopentadienyl-dicarbonylnitrosyl-tungsten) and EDNOW (ethylcyclopentadienyl-dicarbonylnitrosyl-tungsten) may also be used.

In some embodiments, the tungsten-containing gas precursor exposure includes a carrier gas, such as nitrogen (N_2) , argon (Ar), helium (He), hydrogen (H_2) , or other inert gases.

Example pulse times can be between about 0.25 seconds and about 30 seconds, about 0.25 seconds to about 5 seconds, or about 0.5 seconds to about 3 seconds. The pulse may be sufficient to react with the reactive sites on the surface of the feature of the substrate where germanium adsorbed onto the surface.

In some embodiments, both operations 301 and 303 may be performed in a hydrogen environment. After operation 303, there may be an optional purge step to purge excess tungstencontaining precursor in the gas phase.

In operation 305, operations 301 and 303 are repeated until a desired thickness of a tungsten nucleation layer is deposited on the surface of the feature. Each repetition of operations 301 and 303 may be referred to as a "cycle." In some embodiments, method 300 may yield low resistivity tungsten nucleation layers without subsequent treatment. In some embodiments, the order of operations 301 and 303 may be reversed, such that tungsten-containing precursor is pulsed first.

In many embodiments, the deposition of a tungsten nucleation layer will involve pulses of one or more germanium-containing compounds as the only reducing agent pulses, without any pulses of boron-containing reducing agents, silicon-containing reducing agents, or other reducing agents. However, in some embodiments, pulses of one or more additional reducing agents, such as boron-containing or silicon-containing reducing agent pulses, may be used. In these embodiments, the additional reducing agents may be pulsed sequentially or simultaneously with the germanium-containing reducing agent. In this context, it is noted that while hydrogen can be a reducing agent, it is not believed to function as a reducing agent in the nucleation process of FIG. 3 when that is run in a hydrogen environment.

In various embodiments, the amount of tungsten deposited per cycle may be at least about 2 Å, or between about 2 Å and about 8 Å, or less than about 12 Å. As compared to conventional ALD processes, this lower tungsten thickness deposited per cycle allows fine-tuning of the deposition process and overall lower resistivity of the tungsten in the feature. The ability to deposit tungsten nucleation layers with a thickness of between 2 Å and 8 Å allows dialing in of a desire thickness. The tungsten nucleation layer may be deposited to any appropriate thickness. As discussed above, in general, it is advantageous to keep the nucleation layer as thin as possible while still promoting good bulk growth.

As a result, the deposited tungsten nucleation layer may have a thickness less than about 1 nm, or between about 1 nm and about 20 nm, depending on the feature. In various embodiments, the thickness of the tungsten nucleation layer may be between about 1 nm and about 10 nm.

After method 300 is performed, a bulk tungsten layer may be deposited on the tungsten nucleation layer. As an example, the total thickness of the deposited germane-based tungsten nucleation layer and bulk tungsten layer may be between about 1 nm and about 200 nm, or between about 4 nm and 5 about 200 nm, or between about 10 nm and about 50 nm, or between about 1 nm and about 10 nm. Methods of depositing the bulk tungsten film may include CVD, plasma enhanced CVD (PECVD), and physical vapor deposition (PVD). In various embodiments, tungsten grain growth on the nucleation layer deposited using a germanium-containing reducing agent may grow from a first reactive site on the surface of the substrate to the grain growth at a second reactive site on the surface of the substrate. These reactive sites may be where tungsten nucleated during operation 303. In some embodi- 15 ments, these grains may grow horizontally across the width of the feature from one sidewall to another.

Without being bound by a particular theory, it is believed that use of a germanium-containing reducing agent results in formation of metallic, microcrystalline alpha-tungsten 20 (α -W), rather than beta-tungsten (β -W). As discussed above, alpha-tungsten is the stable, crystalline structure of elemental tungsten and has a lower resistivity than beta-tungsten, which is a metastable structure. The formation of the nucleation layer using a germanium-containing reducing agent promotes alpha-tungsten growth during bulk deposition, thereby reducing resistivity overall. The presence of alpha-tungsten versus beta-tungsten may be measured by x-ray diffraction analysis or other suitable methods. The method 300 therefore may be performed such that the tungsten nucleation layer and the 30 subsequent bulk tungsten deposited in the feature is at least 90% alpha-tungsten or at least 99% alpha-tungsten.

An example of a deposited tungsten nucleation layer is shown in FIG. 4A. FIG. 4A is transmission electron microscopy (TEM) image of a deposited tungsten layer in a feature 35 having a 20 nm width. The step coverage for the deposited tungsten was about 85% at the bottom, about 70% on the sidewalls, and about 82% at the opening of the feature. This shows good step coverage for nucleation layers deposited using germanium-containing reducing agents. The deposited using germanium-containing reducing agents. The deposited nucleation layer has a thickness between about 20 Å and about 30 Å. FIG. 4B shows a feature with deposited tungsten bulk layer on a nucleation layer deposited using germane. As shown, the deposited tungsten bulk layer has excellent large grain fill and the nucleation layer itself is very thin and visually indistinguishable from the bulk tungsten layer. This may indicate less of a grain boundary and hence less resistance.

Also the TEM image shows an absence of a seam down the middle of the deposited tungsten film in the feature. This may be due to grains growing from one sidewall of the feature to 50 another. Alternatively, tungsten growth may be extending from one site where tungsten nucleated on the surface of the feature to another site where tungsten nucleated on the surface.

FIG. **5** is a process flow diagram for a method **500** of filling 55 tungsten in a feature of a substrate. As an example, the substrate may be a substrate having features, such as a feature with an aspect ratio of at least 6:1, or at least 10:1, or at least 20:1. In some embodiments, method **500** may be performed at a temperature between about 200° C. and about 400° C. In 60 some embodiments, method **500** may be performed at a pressure between about 1 Torr and about 300 Torr. Temperatures and pressures outside these ranges may be used according to particular implementations.

In operation **501**, a conformal nucleation layer is deposited 65 in the feature. This may be deposited prior to any fill of the feature. This may be performed by any suitable method for

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depositing a nucleation layer, such as ALD, PNL, plasma-enhanced ALD (PEALD), or CVD. In many embodiments, the reducing agent used to deposit the nucleation layer is a germanium-containing reducing agent. In other embodiments, the reducing agent used to deposit the nucleation layer may include boron-containing reducing agents including diborane ($\rm B_2H_6$) and other boranes, silicon-containing reducing agents including silane (SiH₄) and other silanes, hydrazines, and germanes. In some embodiments, the method of depositing a nucleation layer may be a method as described above with respect to FIG. 3.

Next, in operation 503, the nucleation layer is treated by exposing it to pulses of a germanium-containing reducing agent. Examples of a germanium-containing reducing agent include germane (GeH_4) and digermane (Ge_2H_6), and any of those described above with respect to operation 301 in FIG. 3.

In various embodiments, operation 503 may be performed without any intervening pulses of any other precursors. Example pulse times of germanium-containing reducing agent may be between about 0.25 seconds and about 20 seconds, or 0.25 seconds and 5 seconds.

One example of this pulsing scheme without intervening pulses is represented by the schematic illustration in FIG. **6**A. The figure shows an example of a scheme using pulses of germane (GeH₄) over time. As shown, gas flow of each pulse of germane is separated by a time interval. Example time interval pauses between pulses may be between about 0.5 seconds about 5 seconds.

Referring back to FIG. 5, in some embodiments of operation 503, the treatment may be performed with intervening pulses of a tungsten-containing precursor. Examples of tungsten-containing precursors include tungsten hexafluoride (WF₆) and tungsten hexachloride (WCl₆), and any of those described above with respect to operation 303 in FIG. 3.

One example of this pulsing scheme with intervening pulses is represented by the schematic illustration in FIG. 6B. The figure shows an example of the scheme using germane (GeH_4) and tungsten hexafluoride (WF_6), each pulse separated by a time interval.

In some embodiments, the amount of tungsten-containing precursor pulsed during operation 503 in the intervening tungsten-containing precursor pulses may be less than the tungsten-containing precursor pulsed in operation 501. In some embodiments, there may be substantially no tungsten deposited during operation 503 with intervening tungstencontaining precursor pulses, e.g., less than a monolayer is deposited. For example, the pulses of tungsten-containing precursor may be short in duration such that no tungsten adsorbs onto the surface of the tungsten nucleation layer during operation 503. If a germanium-containing reducing agent is used in depositing the nucleation layer, prior to treating it as illustrated in FIG. 6B, transitioning from cycles of the germanium-containing reducing agent/tungsten-containing precursor during nucleation layer deposition to cycles of the germanium-containing reducing agent/tungsten-containing precursor may involve reducing the total amount of tungstencontaining precursor introduced during a pulse, e.g., by shortening a pulse time and/or reducing a flow rate. In various embodiments, operation 503 may be performed in a hydrogen or hydrogen-free environment.

Referring back to FIG. 5, after the nucleation layer is treated, in operation 505, bulk tungsten film is deposited on the tungsten nucleation layer. Methods of depositing the bulk tungsten film may include CVD, PECVD, and PVD.

CVD deposition of a bulk layer can involve flowing a tungsten-containing precursor and a co-reactant such as a reducing agent, if appropriate, into a deposition chamber. An

inert carrier gas may be used to deliver one or more of the reactant streams, which may or may not be pre-mixed. Unlike PNL or ALD processes, this operation generally involves flowing the reactants continuously until the desired amount is deposited. In certain embodiments, the CVD operation may take place in multiple stages, with multiple periods of continuous and simultaneous flow of reactants separated by periods of one or more diverted reactant flows. Flows may also be pulsed for a pulse time between about 1 second and about 2 seconds. Example ranges of chamber pressure during CVD deposition may range from about 10 Torr to about 500 Torr.

Example substrate temperatures are as low as 250° C. and may be as high as 495° C. during the CVD reaction. Various tungsten-containing gases including, but not limited to, WF_6 , $_{15}$ tungsten chloride (WCl₆), and tungsten hexacarbonyl $(W(CO)_6)$, can be used as the tungsten-containing precursor. In certain embodiments, the tungsten-containing precursor is a halogen-containing compound, such as WF₆. In certain embodiments, the reducing agent is hydrogen gas, though 20 other reducing agents may be used, including silane (SiH₄), disilane (Si₂H₆), hydrazine (N₂H₄), diborane (B₂H₆), and germane (GeH₄). In some embodiments, CVD may be implemented in various stages, such as a low temperature stage and a high temperature stage. Still further, nitrogen may be pulsed 25 during CVD deposition of a bulk layer as described in U.S. Pat. No. 8,551,885 and U.S. patent application Ser. No. 13/633,798, both of which are incorporated by reference herein.

FIG. 6C presents a process flow diagram illustrating operations in a method of providing fill according to certain embodiments. The process begins by providing a substrate having a high aspect ratio feature formed therein (602). A germanium-based tungsten nucleation layer is then deposited in the feature as described above, typically to conformally 35 coat the sidewalls and bottom of the feature (604). Examples of nucleation thicknesses range from less than about 1 nm to about 20 nm.

In certain embodiments, depositing the nucleation layer is followed by a post-deposition treatment operation to improve 40 resistivity. Such treatment operations are in U.S. Patent Publication No. 2009/0149022, and U.S. patent application Ser. No. 12/407,541, both of which are incorporated by reference herein, and above with respect to FIGS. **6**A and **6**B. In some embodiments, operations **604** and a post-deposition treatment operation may be boron-free.

Once the nucleation layer is formed, the process continues by filling the feature with a low-temperature CVD tungsten film (606). In this operation, a reducing agent and a tungstencontaining precursor are flowed into a deposition chamber to deposit a bulk fill layer in the feature. An inert carrier gas may be used to deliver one or more of the reactant streams, which may or may not be pre-mixed. Unlike PNL or ALD processes, this operation generally involves flowing the reactants continuously until the desired amount is deposited. In certain 55 embodiments, the CVD operation may take place in multiple stages, with multiple periods of continuous and simultaneous flow of reactants separated by periods of one or more reactant flows diverted.

The reduced temperature CVD operation used to fill the 60 feature can be used with thinner nucleation layers than required by conventional higher temperature CVD. Without being bound by any particular theory, it is believed that this may be because the slower chemistry at the reduced temperatures improves growth even on nucleation sites that are not 65 fully developed. According to various embodiments, nucleation layers of between about 10 Å and about 200 Å may be

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formed; in certain embodiments, nucleation layers may have a thickness less than about 10 Å.

Various tungsten-containing gases including, but not limited to, WF $_6$, WCl $_6$, and W(CO) $_6$ can be used as the tungsten-containing precursor. In certain embodiments, the tungsten-containing precursor is a halogen-containing compound, such as WF $_6$. In certain embodiments, the reducing agent is hydrogen gas, though other reducing agents may be used including silane (SiH $_4$), disilane (Si $_2$ H $_6$) hydrazine (N $_2$ H $_4$), diborane (B $_2$ H $_6$) and germane (GeH $_4$). In many embodiments, hydrogen gas is used as the reducing agent in the CVD process.

CVD filling of the feature is performed at a reduced temperature. According to various embodiments, the reduced temperature (process and/or substrate temperature) is in one of the following ranges: between about 250° C.-350° C., between about 250° C.-340° C., between about 250° C.-325° C., between about 250° C.-325° C., between about 250° C.-315° C., between about 250° C.-310° C., between about 250° C.-315° C., or between about 250° C.-300° C. Also according to various embodiments, the process and/or substrate temperature is: between about 280° C.-310° C., between about 270° C.-310° C., between about 280° C.-310° C., or between about 290° C.-310° C. In certain embodiments, the process and/or substrate temperature is about 300° C.

It should be noted that in some embodiments, a germanium-based nucleation layer may be advantageous for low-temperature CVD fill. This is because, as described above with respect to FIGS. **2**A and **2**B, low-temperature CVD fill on a boron-based nucleation layer may exhibit a rise in resistivity with increasing thickness for certain thicknesses of interest.

After filling the feature, the temperature may be raised to deposit a high temperature CVD layer (608). The high temperature may be in one of the following ranges: between about 350° C.-450° C., between about 360° C.-450° C., between about 370° C.-450° C., between about 380° C.-450° C., between about 390° C.-450° C., or between about 400° C.-450° C. In certain embodiments, the high temperature CVD is performed at about 395° C. Raising the temperature may involve raising the substrate temperature. According to various embodiments, the temperature is raised at least about 50° C., at least about 60° C., at least about 70° C., at least about 80° C., at least about 90° C., at least about 100° C., or at least about 110° C. The high temperature CVD layer is then deposited (610). In certain embodiments, operations 608 and 610 are not performed; that is, after the low temperature CVD process is complete and the feature is filled, the substrate moves on for further processing such as planarization.

In certain embodiments, transitioning from operation 606 to operation 608 involves moving the substrate from one deposition station to another in a multi-station chamber. Still further, each of operation 604, the post-deposition resistivity treatment (if performed), operation 606 and operation 608 is performed in a different station of the same multi-station chamber.

In alternative embodiments in which a single station is used to perform operations 606 and 608, transitioning from operation 606 to operation 608 involved shutting off a flow of tungsten precursor (optionally allowing hydrogen or other reducing gas and/or carrier gas to run), while raising the substrate temperature. Once the substrate temperature is stabilized, the tungsten precursor and other gases, if necessary, are flowed into the reaction chamber for the high temperature deposition. In other embodiments, transitioning from operation 606 may involve raising the substrate temperature while

allowing the deposition to continue during the transition period. In embodiments in which the high temperature tungsten CVD film is deposited, it may be deposited as an overburden layer on the filled feature.

FIG. **6**D illustrates schematic representations of one ⁵ example of a feature's cross-sections at different stages of a filling process in which a high temperature CVD layer is deposited after the feature **650** is filled using reduced temperature CVD. Cross-section **651** represents an example of the feature **650** prior to any tungsten deposition. In this ¹⁰ example, the feature **650** is formed in a dielectric layer **680**, has an opening **675** at the top surface **655** of the substrate and includes a liner layer **653**, such as TiN layer.

APPARATUS

Any suitable chamber may be used to implement the disclosed embodiments. Example deposition apparatuses include various systems, e.g., ALTUS® and ALTUS® Max, available from Lam Research Corp., of Fremont, Calif., or 20 any of a variety of other commercially available processing systems. In some embodiments, the pulsed nucleation process is performed at a first station that is one of two, five or even more deposition stations positioned within a single deposition chamber. Thus, the reducing gases and the tung- 25 sten-containing gases are alternately introduced to the surface of the semiconductor substrate, at the first station, using an individual gas supply system that creates a localized atmosphere at the substrate surface. Another station may be used to for a treatment operation as described above. One or more 30 stations can then be used to perform chemical vapor deposition (CVD) as described above. Two or more stations may be used to perform CVD in a parallel processing. Alternatively a wafer may be indexed to have the CVD operations performed over two or more stations sequentially.

FIG. 7 is a block diagram of a processing system suitable for conducting tungsten thin film deposition processes in accordance with embodiments. The system 700 includes a transfer module 703. The transfer module 703 provides a clean, pressurized environment to minimize risk of contami- 40 nation of substrates being processed as they are moved between various reactor modules. Mounted on the transfer module 703 is a multi-station reactor 709 capable of performing pulsed nucleation layer (PNL) deposition, multi-pulse treatment if desired, and CVD according to embodiments. 45 Chamber 709 may include multiple stations 711, 713, 715, and 717 that may sequentially perform these operations. For example, chamber 709 could be configured such that station 711 performs germane-based PNL deposition, station 713 performs multi-pulse treatment, and stations 715 and 717 can 50 perform CVD. Stations may include a heated pedestal or substrate support, one or more gas inlets or showerhead or

Also mounted on the transfer module 703 may be one or more single or multi-station modules 707 capable of performing plasma or chemical (non-plasma) pre-cleans. The module may also be used for various other treatments, such as a germane treatment. The system 700 also includes one or more wafer source modules 701, where wafers are stored before and after processing. An atmospheric robot (not shown) in the atmospheric transfer chamber 719 may first remove wafers from the source modules 701 to loadlocks 721. A wafer transfer device (generally a robot arm unit) in the transfer module 703 moves the wafers from loadlocks 721 to and among the modules mounted on the transfer module 703.

In various embodiments, a system controller **750** is employed to control process conditions during deposition.

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The controller **750** will typically include one or more memory devices and one or more processors. A processor may include a CPU or computer, analog and/or digital input/output connections, stepper motor controller boards, etc.

The controller **750** may control all of the activities of the deposition apparatus. The system controller **750** executes system control software, including sets of instructions for controlling the timing, mixture of gases, chamber pressure, chamber temperature, wafer temperature, radio frequency (RF) power levels, wafer chuck or pedestal position, and other parameters of a particular process. Other computer programs stored on memory devices associated with the controller **750** may be employed in some embodiments.

Typically there will be a user interface associated with the controller **750**. The user interface may include a display screen, graphical software displays of the apparatus and/or process conditions, and user input devices such as pointing devices, keyboards, touch screens, microphones, etc.

System control logic may be configured in any suitable way. In general, the logic can be designed or configured in hardware and/or software. The instructions for controlling the drive circuitry may be hard coded or provided as software. The instructions may be provided by "programming." Such programming is understood to include logic of any form, including hard coded logic in digital signal processors, application-specific integrated circuits, and other devices which have specific algorithms implemented as hardware. Programming is also understood to include software or firmware instructions that may be executed on a general purpose processor. System control software may be coded in any suitable computer readable programming language.

The computer program code for controlling the germanium-containing reducing agent pulses, hydrogen flow, and tungsten-containing precursor pulses, and other processes in a process sequence can be written in any conventional computer readable programming language: for example, assembly language, C, C++, Pascal, Fortran, or others. Compiled object code or script is executed by the processor to perform the tasks identified in the program. Also as indicated, the program code may be hard coded.

The controller parameters relate to process conditions, such as, for example, process gas composition and flow rates, temperature, pressure, cooling gas pressure, substrate temperature, and chamber wall temperature. These parameters are provided to the user in the form of a recipe, and may be entered utilizing the user interface.

Signals for monitoring the process may be provided by analog and/or digital input connections of the system controller 750. The signals for controlling the process are output on the analog and digital output connections of the deposition apparatus 700.

The system software may be designed or configured in many different ways. For example, various chamber component subroutines or control objects may be written to control operation of the chamber components necessary to carry out the deposition processes in accordance with the disclosed embodiments. Examples of programs or sections of programs for this purpose include substrate positioning code, process gas control code, pressure control code, and heater control code.

A substrate positioning program may include program code for controlling chamber components that are used to load the substrate onto a pedestal or chuck and to control the spacing between the substrate and other parts of the chamber such as a gas inlet and/or target. A process gas control program may include code for controlling gas composition and flow rates and optionally for flowing gas into the chamber

prior to deposition in order to stabilize the pressure in the chamber. A pressure control program may include code for controlling the pressure in the chamber by regulating, e.g., a throttle valve in the exhaust system of the chamber. A heater control program may include code for controlling the current to a heating unit that is used to heat the substrate. Alternatively, the heater control program may control delivery of a heat transfer gas such as helium to the wafer chuck.

Examples of chamber sensors that may be monitored during deposition include mass flow controllers, pressure sensors such as manometers, and thermocouples located in the pedestal or chuck. Appropriately programmed feedback and control algorithms may be used with data from these sensors to maintain desired process conditions.

The foregoing describes implementation of disclosed 15 embodiments in a single or multi-chamber semiconductor processing tool. The apparatus and process described herein may be used in conjunction with lithographic patterning tools or processes, for example, for the fabrication or manufacture of semiconductor devices, displays, LEDs, photovoltaic pan-20 els, and the like. Typically, though not necessarily, such tools/ processes will be used or conducted together in a common fabrication facility. Lithographic patterning of a film typically comprises some or all of the following steps, each step provided with a number of possible tools: (1) application of 25 photoresist on a workpiece, i.e., substrate, using a spin-on or spray-on tool; (2) curing of photoresist using a hot plate or furnace or UV curing tool; (3) exposing the photoresist to visible or UV or x-ray light with a tool such as a wafer stepper; (4) developing the resist so as to selectively remove resist and 30 thereby pattern it using a tool such as a wet bench; (5) transferring the resist pattern into an underlying film or workpiece by using a dry or plasma-assisted etching tool; and (6) removing the resist using a tool such as an RF or microwave plasma resist stripper.

EXPERIMENTAL

Experiment 1

Thickness

Experiments were conducted that show the effect using a germanium-containing reducing agent during deposition of the tungsten nucleation layer had on nucleation layer growth 45 rate and nucleation layer thickness. These experiments were conducted at 300° C.

A first series of experiments involved processing substrates by (1) exposing the substrate to germane (GeH $_4$) without hydrogen present, (2) exposing the substrate to tungsten 50 hexafluoride (WF $_6$), and (3) repeating steps (1) and (2) for various nucleation cycles. Germane was flowed at a flow rate of 250 sccm with a carrier gas argon with a flow rate of 1000 sccm. Tungsten hexafluoride was flowed at a flow rate of 270 sccm. The thickness of the deposited tungsten nucleation 55 layer was measured at various nucleation cycles. The results are shown in FIG. 8 and are represented by the round points.

A second series of experiments involved processing substrates by (1) exposing the substrate to germane (GeH₄) in a hydrogen environment, (2) exposing the substrate to tungsten 60 hexafluoride (WF₆), and (3) repeating steps (1) and (2) for various nucleation cycles. Germane was flowed at a flow rate of 250 sccm with a carrier gas argon with a flow rate of 1000 sccm. Hydrogen was flowed at a flow rate of 19,000 sccm throughout both the germane pulses and the tungsten 65 hexafluoride pulses. Tungsten hexafluoride was flowed at a flow rate of 270 sccm. The thickness of the deposited tungsten

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nucleation layer was measured at various nucleation cycles. The results are represented in FIG. 8 by the square points.

A summary of the tungsten deposition rate (A per cycle) is shown in Table 1.

TABLE 1

	Without 1	Hydrogen	With H	ydrogen
Nucleation Cycle	Thickness (Å)	Thickness per Cycle	Thickness (Å)	Thickness per Cycle
7	4	3.35	23	0.53
9	15	4.27	38	1.62
12	29	5.02	60	2.46
15	42	5.47	82	2.80

As shown, use of hydrogen flow during the tungsten nucleation layer deposition substantially decreased the thickness of the tungsten nucleation layer, which suggests that the presence of hydrogen decreases the amount of tungsten deposited per cycle using germane.

Experiment 2

Resistivity of Nucleation Layers

Experiments were conducted that show the effect using a germanium-containing reducing agent during deposition of the tungsten nucleation layer had on resistivity of the overall deposited tungsten nucleation layer. These experiments were conducted at 300° C.

A first series of experiments involved processing substrates by (1) exposing the substrate to germane (GeH₄) without hydrogen present, (2) exposing the substrate to tungsten hexafluoride (WF₆), and (3) repeating steps (1) and (2) for various nucleation cycles. Germane was flowed at a flow rate of 250 sccm with a carrier gas argon at a flow rate of 1000 sccm. Tungsten hexafluoride was flowed at a flow rate of 270 sccm. The resistivity of the deposited tungsten nucleation layer was measured at various nucleation thicknesses. The results are shown in FIG. 9 and are represented by the round points.

A second series of experiments involved processing substrates by (1) exposing the substrate to germane (GeH₄) in a hydrogen environment, (2) exposing the substrate to tungsten hexafluoride (WF₆), and (3) repeating steps (1) and (2) for various nucleation cycles. Germane was flowed at a flow rate of 250 sccm with carrier gas argon at a flow rate of 1000 sccm. Hydrogen was flowed at a flow rate of 19,000 sccm throughout both the germane (GeH₄) pulses and the tungsten hexafluoride (WF₆) pulses. Tungsten hexafluoride (WF₆) was flowed at a flow rate of 270 sccm. The resistivity of the deposited tungsten nucleation layer was measured at various nucleation thicknesses. The results are represented in FIG. 9 by the square points.

A summary of the resistivity of the nucleation layers deposited using germane is shown in Table 2.

Without Hydrogen		With Hydrogen	
Thickness (Å)	Resistivity $(\mu\Omega\text{-cm})$	Thickness (Å)	Resistivity (μΩ-cm)
15	50.3	23	64.3
29	60.5	38	76.8
42	70.6	60	88.4
		82	98.9

As shown, use of hydrogen flow during the tungsten nucleation layer deposition substantially decreased the resistivity of the tungsten nucleation layer.

The substrate with a tungsten nucleation layer thickness of $20~\mbox{\sc A}$ was further processed and bulk tungsten was deposited into features of the substrate. As a result, the overall resistivity was about $60~\mu\Omega\text{-cm}$, indicating that the resistivity did not change substantially. This is an unexpected result since bulk tungsten layers with thicknesses less than $100~\mbox{\sc A}$ deposited on nucleation layers that were deposited using conventional methods, such as with diborane, generally show a small increase in resistivity after bulk tungsten is deposited. This supports the theory that substantially all of the tungsten deposited into the features may be alpha-tungsten, e.g., low-resistivity tungsten, since $40~\mbox{\sc A}$ of tungsten was deposited over the $20~\mbox{\sc A}$ nucleation layer and the resistivity did not substantially change.

Experiment 3

Overall Resistivity

Experiments were conducted to determine the effect using ³⁵ a germanium-containing reducing agent during deposition of the tungsten nucleation layer had on resistivity of the overall deposited tungsten after bulk tungsten was deposited by chemical vapor deposition (CVD).

Three processes were conducted on substrates to deposit ⁴⁰ nucleation layers and bulk tungsten fill. As an example, process conditions in this experiment are listed in Table 3.

TABLE 3

Process Conditions			
Process	Nucleation Layer	Treatment	Bulk Layer
A	300° C.	None	300° C.
	GeH ₄ /WF ₆ Pulses		CVD
В	300° C.	300° C.	300° C.
	2 cycles of	3 cycles of	CVD
	B2H6/WF6 Pulses	B ₂ H ₆ /WF ₆ Pulses	
С	250° C.	325° ℃.	325° C.
	2 cycles of	5 cycles of	CVD
	B ₂ H ₆ /WF ₆ Pulses	B ₂ H ₆ Pulses	

In Process A, substrates were processed by (1) exposing the substrate to germane (GeH₄) with hydrogen present, (2) exposing the substrate to tungsten hexafluoride (WF₆), (3) repeating steps (1) and (2) for various nucleation cycles, and 60 (4) depositing bulk tungsten by CVD. Substrates were processed at 300° C. for both the nucleation layer deposition and the bulk tungsten deposition. For the nucleation layer deposition, germane was flowed at a flow rate of 250 sccm with a carrier gas argon with a flow rate of 1000 sccm. Hydrogen was 65 flowed at a flow rate of about 19,000 sccm in the front end and 13,000 sccm in the back end. In Processes B and C, substrates

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were processed using diborane (B_2H_6) in two optimized processes. The resistivity of the deposited tungsten layer was measured at various cumulative thicknesses of the nucleation layer and bulk CVD layer. The results are shown in FIG. 10. Process A results are represented by the triangular points, Process B results are represented by the square points, and Process C results are represented by the diamond points. For purposes of this experiment, the point for each process with the lowest thickness is the substrate with only a nucleation layer deposited. Subsequent points corresponding to greater thicknesses represent the resistivities of the overall nucleation layer and deposited tungsten layer by CVD.

Both Process B and Process C involved use of diborane to deposit the nucleation layer, and both of these processes exhibit a sharp increase in resistivity in the overall tungsten film upon deposition of the bulk tungsten layer by CVD. For example, the resistivity of the substrate in Process B increased from 102.6 $\mu\Omega$ -cm to 114.6 $\mu\Omega$ -cm, and the resistivity of the substrate in Process C increased from 83.4 $\mu\Omega$ -cm to 122.1 $\mu\Omega\text{-cm}.$ An unexpected result for Process A is shown. After deposition of the bulk tungsten layer, the resistivity of the substrate decreased from 125.4 $\mu\Omega$ -cm to 108.2 $\mu\Omega$ -cm. According to the trend described above with respect to FIG. 2, this supports the theory that deposition of the nucleation layer using a germanium-containing reducing agent such as germane promotes alpha-tungsten growth because if beta-tungsten were deposited, a sharp increase in resistivity would be shown in the data, similar to the ones represented in Processes 30 B and C. It should be noted that the increase in resistivities for Processes B and C occur at relatively thin thicknesses (e.g., between 40 Å and 80 Å for Process B and about 20 Å and 60 Å for Process C). Accordingly, the germanium-based nucleation process described herein may be particularly suitable for deposition of relatively thin tungsten films, e.g., at less than about 100 Å.

It is noted that although the measurements of resistivity in Process A are higher than some of the resistivities measured in Processes B and C, both Processes B and C have been optimized. Further optimization of the deposition of nucleation layers using germanium-containing compounds for tungsten deposition, decreased overall resistivity may be expected.

Conclusion

Although the foregoing embodiments have been described in some detail for purposes of clarity of understanding, it will be apparent that certain changes and modifications may be practiced within the scope of the appended claims. It should be noted that there are many alternative ways of implementing the processes, systems and apparatus of the present embodiments. Accordingly, the present embodiments are to be considered as illustrative and not restrictive, and the embodiments are not to be limited to the details given herein.

What is claimed is:

- 1. A method of filling a feature on a substrate with tungsten, the method comprising:
 - prior to depositing a bulk tungsten layer, forming a tungsten nucleation layer by exposing the feature to alternating pulses of a germanium-containing reducing agent and a tungsten-containing precursor.
- 2. The method of claim 1, wherein the tungsten nucleation layer is formed to a thickness between 1 nm and 20 nm.
- 3. The method of claim 1, wherein the tungsten nucleation layer is formed to a thickness less than 1 nm.

- **4**. The method of claim **1**, wherein the amount of tungsten deposited per cycle is less than about 8 Å.
- **5**. The method of claim **1**, wherein the tungsten nucleation layer is formed in a hydrogen atmosphere.
- **6**. The method of claim **1**, wherein substantially all of the tungsten formed in the feature comprises alpha-tungsten.
- 7. The method of claim 1, wherein the germanium-containing reducing agent is germane (GeH₄) or digermane (Ge₂H₆).
- **8**. The method of claim **1**, further comprising depositing a $_{10}$ bulk tungsten layer on the tungsten nucleation layer by chemical vapor deposition.
- **9**. The method of claim **8**, wherein grain growth during the bulk tungsten layer deposition extends from a first site where tungsten nucleated to a second site where tungsten nucleated 15 in the feature.
- 10. The method of claim 8, wherein substantially all of the tungsten formed in the feature comprises alpha-tungsten.
- 11. A method of filling a feature on a substrate with tungsten, the method comprising:

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forming a tungsten nucleation layer, and

after forming the tungsten nucleation layer and prior to depositing a tungsten bulk layer, treating the tungsten nucleation layer to pulses of a germanium-containing reducing agent.

12. The method of claim 11, wherein no intervening pulses of other precursors are pulsed during the treatment.

- 13. The method of claim 11, wherein the treatment further comprises intervening pulses of a tungsten-containing precursor, and wherein the amount of the tungsten-containing precursor pulsed during the treatment is less than the amount of the tungsten-containing precursor pulsed during the nucleation layer formation.
- 14. The method of claim 11, wherein the treatment further comprises intervening pulses of a tungsten-containing precursor, wherein substantially no tungsten is deposited during the treatment.
- 15. The method of claim 11, wherein the germanium-containing reducing agent is germane ($Ge_{2}H_{6}$).

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